(19) World Intellectual Property Organization International Bureau



I SENDE DISCUSI IN SONICE MAN COMU COMO RATAL I I IN COMO RECEL MASE ANNO ACESE ANI SOCIALI SECULIA DE PART DEL

(43) International Publication Date 8 January 2004 (08.01.2004)

PCT

(10) International Publication Number WO 2004/003973 A3

- (51) International Patent Classification: *H01J 7/24* (2006.01) *H01J 31/26* (2006.01)
- (21) International Application Number:

PCT/US2003/020197

- (22) International Filing Date: 26 June 2003 (26.06.2003)
- (25) Filing Language: English
- (26) Publication Language: English
- (30) Priority Data: 10/183,768 26 June 2002 (26.06.2002) US 60/463.965 18 April 2003 (18.04.2003) US
- (63) Related by continuation (CON) or continuation-in-part (CIP) to earlier application:

US 10/183,768 (CIP) Filed on 26 June 2002 (26.06.2002)

- (71) Applicant (for all designated States except US): SEME-QUIP INC. [US/US]; 34 Sullivan Road, Unit 21, Billerica, MA 01862 (US).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): HORSKY, Thomas, N. [US/US]; 316 Depot Road, Boxborough, MA 01719 (US). JACOBSON, Dale, C. [US/US]; 16 Flintlock Road, Salem, NH 03079 (US).

- (74) Agents: PANIAGUAS, John, S. et al.; Katten Muchin Zavis Rosenman, Suite 1600, 525 West Monroe Street, Chicago, IL 60661-3693 (US).
- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CC, CC, CC, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, DI, LI, RI, SI, PK, EK, GK, FK, RK, ZL, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, FO, PH, HL, PT, FO, RU, SC, SD, SE, SG, SK, SI, SY, TI, TM, TN, TR, TT, TZ, UA, UG, US, UZ, CV, NY, UZ, AZ, MZ, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, I.S., MW, MZ, SD, SI, SZ, TZ, UG, ZM, ZW), Eumsian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TH), Eumpean patent (AT, BE, BG, CH, CY, CZ, DE, DE, BE, SF, FR, GB, GR, HU, B; TI, LU, MC, NL, PT, RO, SS, SI, SK, TR), OAPT patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

- with international search report
 - before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments
- (88) Date of publication of the international search report: 26 October 2006

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

- (54) Title: ION IMPLANTATION DEVICE AND METHOD
- (57) Abstract: An ion implantation device and a method of manufacturing a semiconductor device is described, wherein ionized boron hydride molecular clusters are implanted to form P-type transitive structures. For example, in the fathreation of Complementary Metal-Oxide Semiconductor (CMOS) devices, the clusters are implanted to provide P-type doping for Source and Drain structures and for Polysters, these objust gets are ceitled to the formation of PMOS transitions. The molecular cluster ions have the chemical form B₂H₂, and B₂H₂, where 10-α>c100 and 0c≤ch+4. The use of such boron hydride clusters results in a dramatic increase in water throughput, as well as improved device yields through the reduction of wafer charging. Thus, this technology significantly reduces manufacturing costs relative to prior implantation techniques. A method of manufacturing a semiconductor device is further described, comprising the steps of providing a supply of molecules containing a plurality of dopant atoms into an ionization chamber, ionizing said molecules into dopant cluster ions, extracting and accelerating the depart cluster ions with an electric field, selecting the desired cluster ions by mass analysis, modifying the final implant energy of the cluster for through post-analysis ion optics, and implanting the dopant cluster ions into an ionization charge, which is an integer number greater than 10. This method enables increasing the dopant doser rate to n times the implantation current with an equivalent per dopant atoms energy of I/m times the cluster implantation energy, while reducing the charge per dopant atom by the factor n. This is an effective method for making shallow transistor junctions, where it is desired to implant with a low energy per dopant atom.

International application No.

PCT/US03/20197

A. CLASSIFICATION OF SUBJECT MATTER						
IPC(7) : H01J 7/24, 31/26						
	US CL : 250/427, 424, 425, 423R; 315/111.81, 111.91					
	International Patent Classification (IPC) or to both n	ational classification and IPC				
	DS SEARCHED					
	Minimum documentation searched (classification system followed by classification symbols) U.S.: 250/281, 427, 424, 425, 423R; 315/111.81, 111.91; 437/165; 427/530; 204/192P;					
Documentati	Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched					
Electronic da	ata base consulted during the international search (nan	ne of data base and, where practicable, sea	rch terms used)			
C. DOC	UMENTS CONSIDERED TO BE RELEVANT					
Category *	Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to claim No.			
Y	US 5,489,550 (MOSLEHI) 06 February 1996 (06.0		1, 5, 6, 10-12, 15, 16			
	column 2, lines 53-55	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	1,0,0,1012,10,10			
Y	US 5,993,766 (TOM et al.) 30 November 1999 (30 14, lines 9-15	1.11.1999), col.9, lines 43-45 and column	1, 5, 6, 10-12, 15, 16			
Y	US 6,013,332 (GOTO et al.) 11 January 2000 (11.0	01.2000), column 2, lines 39-58	1, 5, 6, 10-12, 15, 16			
Y	US 4,412,900 (TANAKA et al.) 01 November 198	3 (01.11.1983), column 2, lines 32-38	1, 5, 6, 10, 15, 16			
Y	US 4,412,900 (TANAKA et al.) 01 November 198	3 (01.11.1983), column 2, lines 32-38	11, 12			
Y	US 4,412,900 (TANAKA et al.) 01 November 198 and 55-56	3 (01.11.1983), column 12, lines.46-48	6, 16			
P	US 2003/0030010 A1 (PEREL et al.) 13 February	2003 (13.02.2003), section [0021]	5, 15			
P	US 2003/0111014 A1 (DONATUCCI et al.) 19 Jun	e 2003 (19.06.2003), section [0038]	5, 15			
	documents are listed in the continuation of Box C.	See patent family annex.				
* S	pecial categories of cited documents:	"T" later document published after the inter	mational filing date or priority			
	defining the general state of the art which is not considered to be lar relevance	date and not in conflict with the applica- principle or theory underlying the lave	ation			
	"X" document of particular relevance: the claimed invention cannot be		claimed invention cannot be red to involve an inventive step			
"L" document establish t specified)	which may throw doubts on priority claim(s) or which is cited to the publication date of another citation or other special reason (as	"Y" document of particular relevance; the considered to lovelye an inventive step	when the document is			
"O" document referring to an oral disclosure, use, exhibition or other means being obvious to a person skilled in the art		documents, such combination				
"P" document priority di	"P" document published prior to the internstional filing date but later than the priority date chilmed					
	cual completion of the international search	Date of mailing of the international searc	h report			
29 September 2004 (29.09.2004)						
Name and mailing address of the ISA/US Mail Stop PCT, Atm: ISA/US		Authorized officer Phonolog for	_ Bell			
Corr	missioner for Patents	John R Lee	-			
P.O.	P.O. Box 1450 Alexandria, Virginia 22313-1450 Telephone No. 703 308 0956					
	Facsimile No. (703)305-3230					
Form PCT/ISA	Form PCT/ISA/210 (second sheet) (July 1998)					

PCT/US03/20197

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
T T	US 6,744.214 B2 (HORSKY) 01 June 2004 (01.06.2004), column 8, lines 1-5	28-30
	US 6.452,338 B1 (HORSKY) 17 September 2002 (17.09.2002), cloumn 7, lines 61-67	28-30
T, P		
Y	US 5,313,061 (DREW et al.) 17 May 1994 (17.05.1994), column 3, lines 63-67	28-30
Y	US 4,587,432 (AITKEN) 06 May 1986 (06.05.1986), column 7, lines 28-36	28-30
Y	US 4,587,432 (AITKEN) 06 May 1986 (06.05.1986), Abstract, column 11, lines 48-58 and lines 66-68, column 12, lines 3-5	29
Y	US 5,313,061,{DREW et al.) 17 May 1994 (17.05.1994), column 11 lines 45-48, column 12, lines 3-10 and column 16, lines 1-9	28
T	US 6,744,214 B2 (HORSKY) 01 June 2004 (01.06.2004), column 12, lines 3-21, column 12, lines 33-35 and column 6, line 46 and lines 33-48	28-30, 34, 36-39
T	US 6,452,338 BI (HORSKY) 17 September 2002 (17.09.2002), column 5, lines 60-65 and Col.6, lines 33-48	28-30, 34, 36-39
	7	

International application No.
PCT/US03/20197

Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)		
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:		
Claim Nos.: because they relate to subject matter not required to be searched by this Authority, namely:		
Claim Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:		
Claim Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(e).		
Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)		
This International Searching Authority found multiple inventions in this international application, as follows: Please See Continuation Sheet		
As all required additional search fees were timely paid by the applicant, this international search report covers all		
searchable claims. 2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.		
As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:		
No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.		

Form PCT/ISA/210 (continuation of first sheet(1)) (July 1998)

PCT/US03/20197	-

Continuation of Item 4 of the first sheet: The tile is too long. NEW TITLE: "fon Implanation Device and Method"
BOX II. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING This application contains the following inventions or groups of inventions which are not so linked as to form a single general inventive concept under PCT Rule 131. However, all claims could be searched without effort justifying an additional fee. Group I, claim(s) 1-19, 24-27 and 28-39, drawn to ion implantation and ion source used for producing implant-ions. Group II, claim(s) 2023, drawn to magnetic yoke assembly that could —but not must— be used in claims of Group I.
·

Form PCT/ISA/210 (second sheet) (July 1998)

(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 30 June 2005 (30.06.2005)

(10) International Publication Number WO 2005/059942 A2

(51) International Patent Classification7:

PCT TIO11

- (21) International Application Number:

PCT/US2004/041525

- (74) Agent: WILLIAMS, John, N.; Fish & Richardson P.C., 225 Franklin Street, Boston, MA 02110-2804 (US).
- (22) International Filing Date: 9 December 2004 (09.12.2004)
- (25) Filing Language:

English

(26) Publication Language:

English

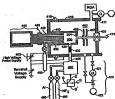
- (30) Priority Data: 60/529,343
 - 12 December 2003 (12.12.2003) US
- (71) Applicant (for all designated States except US): SEME-QUIP, INC. [US/US]; 34 Sullivan Road, Unit 21, Billerica, MA 01862 (US).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): HORSKY, Thomas, N. [US/US]; 816 Depot Road, Boxborough, MA 01719 (US). MILGATE, Robert, W., III [-/US]; 2 Heath Heights, Gloucester, MA 01930 (US), SACCO, George, P., Jr. [-/US]; 51 Parsonage Lane, Topsfield, MA 01983 (US). JACOBSON, Dale, Conrad [-/US]; 16 Flintlock Road, Salem, NH 03079 (US). KRULL, Wade, Allen [-/US]; 8 Smith Street, Marblehead, MA 01945 (US).
- (81) Designated States (unless otherwise indicated, for every kind of national protection available): AB, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, BC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM,
- (84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), Buropean (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SL, SK, TR), OAPI (BF, BJ, CF, CG, CL, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

without international search report and to be republished upon receipt of that report

[Continued on next page]

(54) Title: METHOD AND APPARATUS FOR EXTENDING EQUIPMENT UPTIME IN ION IMPLANTATION



(57) Abstract: The service lifetime of an ion source is enhanced or prolonged by the source having provisions for in-situ etch cleaning of the ion source and of an extraction electrode, using reactive halogen gases, and by having features that extend the service duration between cleanings. The latter include accurate vapor flow control, accurate focusing of the ion beam optics, and thermal control of the extraction electrode that prevents formation of deposits or prevents electrode destruction. An apparatus comprises of an ion source for generating dopant ions for semiconductor wafer processing is coupled to a remote plasma source which delivers For Cl ions to the first ion source for the purpose of cleaning deposits in the first ion source and the extraction electrode. These methods and apparatus enable long equipment uptime when running condensable feed gases such as sublimated vapor sources, and are particularly applicable for use with so-called cold ion sources. Methods and apparatus are described which enable long equipment uptime when decaborane and octadecarborane are used as feed materials, as well as when vaporized elemental arsenic and phosphorus are used, and which serve to enhance beam stability during ion implantation.

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

METHOD AND APPARATUS FOR EXTENDING EQUIPMENT UPTIME IN ION IMPLANTATION

FIELD OF THE INVENTION

The present invention relates to producing ion beams in which one or more gaseous or vaporized feed materials is ionized in an ion source. It also relates to a method and apparatus for operating an ion source to produce an ion beam for ion implantation of semiconductor substrates and substrates for flat panel displays. In particular the invention concerns extension of the productive time (i.e. the "uptime") of systems that produce ion beams.

BACKGROUND

Ion beams are produced from ions extracted from an ion source. An ion source typically employs an ionization chamber connected to a high voltage power supply. The ionization chamber is associated with a source of ionizing energy, such as an arc discharge, energetic electrons from an electron-emitting cathode, or a radio frequency or microwave antenna, for example. A source of desired ion species is introduced into the ionization chamber as a feed material in gaseous or vaporized form where it is exposed to the ionizing energy. Extraction of resultant ions from the chamber through an extraction aperture is based on the electric charge of the ions. An extraction electrode is situated outside of the ionization chamber, aligned with the extraction aperture, and at a voltage below that of the ionization chamber. The electrode draws the ions out, typically forming an ion beam. Depending upon desired use, the beam of ions may be mass-analyzed for establishing mass and energy purity, accelerated, focused and subjected to scanning forces. The beam is then transported to its point of use, for example into a processing chamber. As the result of the precise energy qualities of the ion beam, its ions may be implanted with high accuracy at desired depth into semiconductor substrates.

The precise qualities of the ion beam can be severely affected by condensation and deposit of the feed material or of its decomposition products on surfaces of the ion beam-producing system, and in particular surfaces that affect ionization, ion extraction and acceleration.

The Ion Implantation Process

The conventional method of introducing a dopant element into a semiconductor wafer is by introduction of a controlled energy ion beam for ion implantation. This introduces desired impurity species into the material of the semiconductor substrate to form doped (or "impurity") regions at desired depth. The impurity elements are selected to bond with the semiconductor material to create electrical carriers, thus altering the electrical conductivity of the semiconductor material. The electrical carriers can either be electrons (generated by N-type dopants) or "holes" (i.e., the absence of an electron), generated by P-type dopants. The concentration of dopant impurities so introduced determines the electrical conductivity of the doped region. Many such N- and P-type impurity regions must be created to form transistor structures, isolation structures and other such electronic structures, which collectively function as a semiconductor device.

To produce an ion beam for ion implantation, a gas or vapor feed material is selected to contain the desired dopant element. The gas or vapor is introduced into the evacuated high voltage ionization chamber while energy is introduced to ionize it. This creates ions which contain the dopant element (for example, in silicon the elements As, P, and Sb are donors or N-type dopants, while B and In are acceptors or P-type dopants). An accelerating electric field is provided by the extraction electrode to extract and accelerate the typically positively-charged ions out of the ionization chamber, creating the desired ion beam. When high purity is required, the beam is transported through mass analysis to select the species to be implanted, as is known in the art. The ion beam is ultimately transported to a processing chamber for implantation into the semiconductor wafer.

Similar technology is used in the fabrication of flat-panel displays (FPD's) which incorporate on-substrate driver circuitry to operate the thin-film transistors which populate the displays. The substrate in this case is a transparent panel such as glass to which a semiconductor layer has been applied. Ion sources used in the manufacturing of FPD's are typically physically large, to create large-area ion beams of boron, phosphorus and arsenic-containing materials, for example, which are directed into a chamber containing the substrate to be implanted. Most FPD implanters do not mass-analyze the ion beam prior to its reaching the substrate.

Ion Contamination

In general, ion beams of N-type dopants such as P or As should not contain any significant portion of P-type dopant ions, and ion beams of P-type dopants such as B or In should not contain any significant portion of N-type dopant ions. Such a condition is called "cross-contamination" and is undesirable. Cross-contamination can occur when source feed materials accumulate in the ion source, and the source feed material is then changed, for example, when first running elemental phosphorus feed material to generate an N-type P' beam, and then switching to BF₃ gas to generate a P-type BF₂⁺ beam.

A serious contamination effect occurs when feed materials accumulate within the ion source so that they interfere with the successful operation of the source. Such a condition invariably has called for removal of the ion source and the extraction electrode for cleaning or replacement, resulting in an extended "down" time of the entire ion implantation system, and consequent loss of productivity.

Many ion sources used in ion implanters for device wafer manufacturing are "hot" sources, that is, they operate by sustaining an arc discharge and generating a dense plasma; the ionization chamber of such a "hot" source can reach an operating temperature of 800C or higher, in many cases substantially reducing the accumulation of solid deposits. In addition, the use of BF₃ in such sources to generate boron-containing ion beams further reduces deposits, since in the generation of a BF₃ plasma, copious amounts of fluorine ions are generated; fluorine can etch the walls of the ion source, and in particular, recover deposited boron through the chemical production of gaseous BF₃. With other feed materials, however, detrimental deposits have formed in hot ion sources. Examples include antimony (Sb) metal, and solid indium (In), the ions of which are used for doping silicon substrates.

Cold ion sources, for example the RF bucket-type ion source which uses an immersed RF antenna to excite the source plasma (see, for example, Leung et al., U.S. Patent 6,094,012, herein incorporated by reference), are used in applications where either the design of the ion source includes permanent magnets which must be kept below their Curie temperature, or the ion source is designed to use thermally-sensitive feed materials which break down if exposed to hot surfaces, or where both of these conditions exist. Cold ion sources suffer more from the deposition of feed materials than do hot sources.

The use of halogenated feed materials for producing dopants may help reduce deposits to some extent, however, in certain cases, non-halogen feed materials such as hydrides are preferred over halogenated compounds. For non-halogen applications, ion source feed materials such as gaseous B₂H₆, AsH₃, and PH₃ are used. In some cases, elemental As and P are used, in vaporized form. The use of these gases and vapors in cold ion sources has resulted in significant materials deposition and has required the ion source to be removed and cleaned, sometimes frequently. Cold ion sources which use B₂H₆ and PH₃ are in common use today in FPD implantation tools, These ion sources suffer from cross-contamination (between N- and P-type dopants) and also from particle formation due to the presence of deposits. When transported to the substrate, particles negatively impact yield. Cross-contamination effects have historically forced FPD manufacturers to use dedicated ion implanters, one for N-type ions, and one for P-type ions, which has severely affected cost of ownership.

Borohydrides

Borohydride materials such as $B_{10}H_{14}$ (decaborane) and $B_{18}H_{22}$ (octadecaborane) have attracted interest as ion implantation source materials. Under the right conditions, these materials form the ions $B_{10}H_x^+$, $B_{10}H_x^-$, $B_{18}H_x^+$, and $B_{18}H_x^-$. When implanted, these ions enable very shallow, high dose P-type implants for shallow junction formation in CMOS manufacturing. Since these materials are solid at room temperature, they must be vaporized and the vapor introduced into the ion source for ionization. They are low-temperature materials (e.g., decaborane melts at 100C, and has a vapor pressure of approximately 0.2 Torr at room temperature; also, decaborane dissociates above 350C), and hence must be used in a cold ion source. They are fragile molecules which are easily dissociated, for example, in hot plasma sources.

Contamination Issues of Borohydrides

Boron hydrides such as decaborane and octadecaborane present a severe deposition problem when used to produce ion beams, due to their propensity for readily dissociating within the ion source. Use of these materials in Bernas-style are discharge ion sources and also in electron-impact ("soft") ionization sources, have confirmed that

boron-containing deposits accumulate within the ion sources at a substantial rate. Indeed, up to half of the borohydride vapor introduced into the source may stay in the ion source as dissociated, condensed material. Eventually, depending on the design of the ion source, the buildup of condensed material interferes with the operation of the source and necessitates removal and cleaning of the ion source.

Contamination of the extraction electrode has also been a problem when using these materials. Both direct ion beam strike and condensed vapor can form layers that degrade operation of the ion beam formation optics, since these boron-containing layers appear to be electrically insulating. Once an electrically insulating layer is deposited, it accumulates electrical charge and creates vacuum discharges, or so-called "gittches", upon breakdown. Such instabilities affect the precision quality of the ion beam and can contribute to the creation of contaminating particles.

SUMMARY OF THE INVENTION

It is an object of this invention to provide a method and apparatus for producing an ion beam which increases service lifetime and reduces equipment down time.

The invention features in-situ cleaning procedures and apparatus for an ion source and associated extraction electrodes and similar components of the ion-beam producing system, which periodically chemically remove deposits, increasing service lifetime and performance, without the need to disassemble the system.

The invention also features an actively heated ion extraction electrode which consists of a material which reduces the frequency and occurrence of electrical discharges, preferably this material being a metal.

Another feature is, in general, heating an extraction electrode above the condensation temperature of feed materials to an ion source, in preferred cases the electrode being comprised of metal, preferably aluminum or molybdenum.

The invention also features steps and construction features that improve the production and utilization of ions. This reduces the required concentration of feed material in the ion source, and in related manner, reduces the rate of accumulation of detrimental deposits on surfaces of the ion-beam producing system.

These and other innovations of the invention may include one or more of the following features:

A supply of a reactive gas is provided and introduced into the ion source, and the ion source is cleaned in sim through exposure to reactive products from that supply such as atomic fluorine, F, or molecular fluorine, F₂; the atomic or molecular fluorine is injected into the ion source from a remote plasma source; the ion source is cleaned through exposure to gaseous CIF₃ flowing from a remote supply; reactive components of the ionization apparatus are shielded from reactive gas during the cleaning phase of operation; the ion source is fabricated of aluminum; the extraction electrode is fabricated of aluminum; the front face of the extraction electrode is devoid of sharp or rough features; the plates of the extraction electrode are actively temperature controlled; the plates of the extraction electrode are actively heated; heating of the extraction electrode is actively or is resistive; the plates of the extraction electrode in other situations are actively cooled.

Another feature is the use of the features described with apparatus suitable to form "cluster" or "molecular" ion beams, of feed material that is particularly subject to thermal breakdown and deposit. . . .

While most ion implantation experts would agree that the use of borohydrides to form "cluster" ion beams such as $B_{10}H_{\star}^{**}$ and $B_{10}H_{\star}^{**}$ is very attractive for shallow junction formation, means to ionize and transport these large molecules have presented problems. For example, U.S. patents 6,288,403 and 6,452,338 describe ion sources which have successfully produced decaborane ion beams. However, such decaborane ion sources have been found to exhibit particularly short service life as compared to other commercial ion sources used in ion implantation. This short service life has been primarily due to the accumulation of boron-containing deposits within the ion source, and the deposition of insulating coatings on the ion extraction electrode, which has lead to beam instabilities requiring implanter shut down and maintenance.

According to another feature, means are provided to substantially reduce the deposition of such deposits in the borohydride ion source and on the ion extraction electrode, and means are provided to clean deposits on these components without removing them from the ion implanter, i.e., in-situ. This invention enables the

commercial use of borohydride cluster beams in semiconductor manufacturing with long service lifetime.

A particular aspect of the invention is a system for generating an ion beam comprising an ion source in combination with an extraction electrode and a reactive gas cleaning system, the ion source comprising an ionization chamber connected to a high voltage power supply and having an inlet for gaseous or vaporized feed materials, an energizeable ionizing system for ionizing the feed material within the ionization chamber and an extraction aperture that communicates with a vacuum housing, the vacuum housing evacuated by a vacuum pumping system, the extraction electrode disposed in the vacuum housing outside of the ionization chamber, aligned with the extraction aperture of the ionization chamber and adapted to be maintained at a voltage below that of the ionization chamber to extract ions through the aperture from within the ionization chamber, and the reactive gas cleaning system operable when the ionization chamber and ionizing system are de-energized to provide a flow of reactive gas through the ionization chamber and through the ion extraction aperture to react with and remove deposits on at least some of the surfaces of the ion generating system.

Preferred embodiments of this aspect have one or more of the following features.

The system is constructed for use in implanting ions in semiconductor wafers, the ionization chamber having a volume less than about 100ml and an internal surface area of less than about 200cm².

The system is constructed to produce a flow of the reactive gas into the ionization chamber at a flow rate of less than about 2 Standard Liters Per Minute.

The extraction electrode is constructed to produce a beam of accelerated ions suitable for transport to a point of utilization.

The extraction electrode is located within a path of reactive gas moving from the extraction aperture to the vacuum pumping system so that the extraction electrode is cleaned by the reactive gas.

The extraction electrode is associated with a heater to maintain the electrode at elevated temperature during extraction by the extraction electrode of ions produced in the

ionization chamber, e.g. above the condensation temperature, below the disassociation temperature, of solid-derived, thermally sensitive vapors.

The extraction electrode is associated with a cooling device, e.g. when the electrode is formed of thermally sensitive material and is used with a hot ion source.

The extraction electrode has a smooth, featureless aspect.

The reactive gas cleaning system comprises a plasma chamber, the plasma chamber arranged to receive a feed gas capable of being disassociated by plasma to produce a flow of reactive gas through a chamber outlet, and a conduit for transporting the reactive gas to the ionization chamber.

The plasma chamber is constructed and arranged to receive and disassociate a compound capable of being disassociated to atomic fluorine, for instance NF₃, C_3F_8 or CF4.

The reactive gas cleaning system is constructed and arranged to share a service facility associated with the ion source.

The system is constructed to direct an ion beam through a mass analyzer, in which the reactive gas cleaning system is constructed and arranged to share a service facility with the mass analyzer.

The reactive gas cleaning system comprises a conduit from a container of pressurized reactive gas, for instance CIF₃.

The system is in combination with an end-point detection system adapted to at least assist in detecting substantial completion of reaction of the reactive gas with contamination on a surface of the system for generating an ion beam.

The end point detection system comprises an analysis system for the chemical makeup of gas that has been exposed to the surface during operation of the reactive gas cleaning system.

A temperature detector is arranged to detect substantial termination of an exothermic reaction of the reactive gas with contamination on a surface of the system.

The energizeable ionizing system includes a component within or in communication with the ionization chamber that is susceptible to harm by the reactive gas and means are provided to shield the component from reactive gas flowing through the system.

The means to shield the component comprises an arrangement for producing a flow of inert gas, such as argon, past the component.

The means for shielding a component comprises a shield member that is impermeable to the reactive gas.

The system is constructed to operate with reactive halogen gas as the reactive gas and the extraction electrode and associated parts comprise aluminum (A1) or alumina (AlOs).

The ion source is constructed to produce ions within the ionization chamber via an arc-discharge, an RF field, a microwave field or an electron beam.

The system is associated with a vaporizer of condensable solid feed material for producing feed vapor to the ionization chamber.

The ion source is constructed to vaporize feed material capable of producing cluster or molecular ions, and the ionization system is constructed to ionize the material to form cluster or molecular ions for implantation.

The vacuum housing of the system is associated with a pumping system comprising a high vacuum pump capable of producing high vacuum and a backing pump capable of producing vacuum, the high vacuum pump operable during operation of the ion source, and being capable of being isolated from the vacuum housing during operation of the reactive cleaning system, the backing pump operable during operation of the reactive gas cleaning system.

The system is associated with an ion implantation apparatus, the apparatus constructed to transport ions following the extraction electrode implantation station within a vacuum chamber. In preferred embodiments an isolation valve is included for isolating the implantation station from the ionization chamber and the extraction electrode during operation of the reactive gas cleaning system.

The ion source is constructed and adapted to generate dopant ions for semiconductor processing, and the reactive gas cleaning system is adapted to deliver fluorine, F, or chlorine, Cl, ions to the ionization chamber or the extraction electrode for cleaning deposits from a surface.

The ion source is adapted to be temperature-controlled to a given temperature.

The ion source is adapted to generate a boron-containing ion beam; in preferred embodiments the boron-containing ion beam is generated by feeding vaporized borohydride material into the ion source, especially either decaborane, $B_{10}H_{14}$ or octadecaborane, $B_{18}H_{22}$.

The ion source is adapted to generate arsenic-containing ion beams.

The ion source is adapted to generate phosphorus-containing ion beams.

The ionization chamber of the ion source comprises aluminum.

The ionization chamber of the ion source or the extraction electrode comprises a material resistant to attack by halogen gases such as fluorine, F.

Another particular aspect of the invention is a method of *in-situ* cleaning using the system of any of the foregoing description, or of an ion source associated with an ion implanter, in which reactive halogen gas is flowed into an ion source while the ion source is de-energized and under vacuum.

Embodiments of this aspect have one or more of the following features.

The reactive halogen gas is fluorine, F.

The reactive halogen gas is chlorine, Cl.

The fluorine gas is introduced into the ion source from a remote plasma source.

The fluorine gas is produced in the remote plasma source by an NF3 plasma.

The fluorine gas is produced in the remote plasma source by a C_3F_8 or CF_4 plasma.

The reactive halogen gas is CIF₃.

The cleaning procedure is conducted to remove deposits after the ion source has ionized decaborane, B $_{10}$ H $_{14}$.

The cleaning procedure is conducted to remove deposits after the ion source has ionized octadecaborane, B₁₈H₂₂.

The cleaning procedure is conducted to remove deposits after the ion source has ionized arsenic-containing compounds, such as arsine, AsH₃, or elemental arsenic, As.

The cleaning procedure is conducted to remove deposits after the ion source has ionized phosphorus-containing compounds, such as elemental phosphorus, P, or phosphine, PH₃.

The cleaning procedure is conducted to remove deposits after the ion source has ionized antimony-containing compounds, such as trimethylantimony, Sb(CH₄)₅, or antimony pentaflouride, SbF₅.

The cleaning procedure is conducted for an ion source in situ in an ion implanter between changing ion source feed materials in order to implant a different ion species.

Another particular aspect of the invention is an ion implantation system having an ion source and an extraction electrode for extracting ions from the ion source, in which the extraction electrode includes a heater constructed to maintain the electrode at an elevated temperature sufficient to substantially reduce condensation on the electrode of gases or vapors being ionized and products produced therefrom.

Embodiments of this aspect have one or more of the following features.

· The electrode comprises aluminum.

The electrode comprises of molybdenum.

The electrode is heated by radiative heaters.

The electrode is heated by resistive heaters).

The temperature of the electrode is controlled to a desired temperature; in embodiments the temperature is between 150C and 250C.

The electrode is periodically cleaned in situ by exposure to reactive halogencontaining gas.

Another aspect of the invention is a method of *in-situ* cleaning of an ion extraction electrode of any of the systems described or an ion extraction electrode which is associated with an ion implanter, in which reactive halogen gas is flowed over the ion extraction electrode while the electrode is *in situ* and under vacuum.

Embodiments of this aspect have one ore more of the following features.

The reactive halogen gas is fluorine, F or chlorine, Cl.

Fluorine gas is introduced from a remote plasma source into a vacuum housing in which the extraction electrode resides.

Fluorine gas is produced in the remote plasma source by a NF3 plasma.

Fluorine gas is produced in the remote plasma source by a C₃F₈ or CF₄ plasma.

The reactive gas is CIF₃.

The cleaning procedure is conducted to remove deposits after the ion source has ionized decaborane, $B_{10}H_{14}$.

The cleaning procedure is conducted to remove deposits after the ion source has ionized octadecaborane, B₁₈H₂₂.

The cleaning procedure is conducted to remove deposits after the ion source has ionized arsenic-containing compounds, such as arsine, AsH₃, or elemental arsenic, As.

The cleaning procedure is conducted to remove deposits after the ion source has ionized phosphorus-containing compounds, such as elemental phosphorus, P, or phosphine, PH₃.

The cleaning procedure is conducted between changing ion source feed materials in order to implant a different ion species.

Ion Source Provided With In-situ Etch Cleaning

According to a preferred embodiment, the *in situ* chemical cleaning process utilizes atomic F gas, to effectively clean deposits from the ion source and from the ion extraction electrode, while the ion source and extraction electrode remain installed in the ion beam-producing system. In a preferred embodiment an electron impact ion source with cooled chamber walls is employed. Preferably, the ionization chamber and source block are fabricated of aluminum or of an aluminum containing complex, enabling aluminum fluoride to be created on the aluminum surfaces to act as a passivating layer, that prevents further chemical attack by F.

One embodiment of this feature uses the outlet of a remote reactive gas source directly coupled to an inlet to the ion source.

In a preferred embodiment the reactive gas source is a plasma source which introduces an etch feed gas, such as $N\Gamma_2$ or $C_3\Gamma_8$, into a supplemental ionization chamber. By sustaining a plasma in the supplemental chamber, reactive gases such as F and Γ_2 are produced, and these reactive gases, introduced to the main ion source, chemically attack the deposited materials. By-products released in the gas phase are drawn through the extraction aperture of the ionization chamber and are pumped away by the vacuum system of the installation, cleaning the chamber.

Deposition Model

It is a generally observed principle of physics that when two objects interact, there can be more than one outcome. Furthermore, one can assign probabilities or likelihoods to each outcome such that, when all possible outcomes are considered, the sum of their individual probabilities is 100%. In atomic and molecular physics such possible outcomes are sometimes called "channels" and the probability associated with each interaction channel is called a "cross section". More precisely, the likelihood of two particles (say, an electron and a gas molecule) interacting with each other at all is the "total cross section", while the likelihoods of certain types of interactions (such as the interaction represented by the electron attaching itself to the gas molecule thus forming a negative ion, or by removing an electron from the gas molecule thus forming a positive ion, or by dissociating the molecule into fragments, or by elastically scattering from the molecule with no chemical change of the molecule) are the "partial cross sections".

This state of affairs can be represented by a mathematical relation which expresses the total cross section σ_T as the sum of its i partial cross sections:

(1)
$$\sigma_T = \sigma_1 + \sigma_2 + \sigma_3 + \dots + \sigma_i$$
, or

(2)
$$\sigma_T = \sum \sigma_i$$
.

The ion sources used in ion implanters typically display modest ionization fractions. That is, only a small fraction (from a few per cent to a few tens of per cent) of the gas or vapor fed into the ion source is ionized. The rest of the gas or vapor typically leaves the source in the gas phase, either in its original state or in some other neutral state. That is, the ionization cross section is much smaller than the total cross section. Of course, some of the gas components can stay in the ion source as deposited materials, although this tends to be a small percentage of the total for the commonly used implantation feed materials. While feed materials vaporized by heating such as elemental As or P more readily produce deposits than do normally gaseous feed materials, the heated vapor tends to stay in the gas phase if the walls of the ion source are at a higher temperature than the vaporizer, and do not pose a severe deposition risk. However, significant detrimental deposits may still be produced when producing boron beams from gaseous BF3 feed gas, for example, as well as beam from In and Sb.

Also, in general, over time, deposits of condensable materials do occur on certain other components of ion producing systems, affecting their operational life before disassembly and cleaning.

Furthermore, in the case of the borohydrides, the total cross section representing all interactions with the ionizing medium (i.e., electrons in the ion source) appears large, the ionization cross section is small, and by far the largest cross section represents the channel for dissociation of the borohydride molecules into non-volatile fragments, which then remain on surfaces in the ion source. The problem of deposition of these fragments is adversely influenced by cooling of the ionization chamber walls in an effort to reduce thermal decomposition of the feed material. In sum, it appears that deposition from borohydrides of boron-containing fragments in the source is a fundamental phenomenon which would be observed in any type of ion source acting on this material, and solution to the problem is of broad, critical interest to the semiconductor manufacturing industry.

Electron Impact Ion Source Suitable For Borohydrides

An ion source particularly suitable for borohydrides is an electron-impact ion source which is fully temperature-controlled (see U.S. Patents 6,452,338 and 6,686,595; also International Application no. PCT/US03/20197, each herein incorporated by reference); also see FIG. 7. Instead of striking an arc-discharge plasma to create ions, the ion source uses a "soft" electron-impact ionization of the process gas by energetic electrons injected in the form of one or more focused electron beams. This "soft" ionization process preserves these large molecules so that ionized clusters are formed. As : seen in FIG. 7, solid borohydride material is heated in a vaporizer and the vapor flows through a vapor conduit to a metal chamber, i.e., the ionization chamber. An electron gun located external to the ionization chamber delivers a high-current stream of energetic electrons into the ionization chamber; this electron stream is directed roughly parallel and adjacent to an extended slot in the front of the chamber. Ions are extracted from this slot by an ion extraction electrode, forming an energetic ion beam. During transport of the sublimated borohydride vapor to the ionization chamber all surfaces are held at a higher temperature than that of the vaporizer (but well below the temperature of dissociation), to prevent condensation of the vapor. Many hours of testing have confirmed that the

surfaces of the vapor feed and valves indeed remain clean when such temperature control is implemented.

Extension of Ion Source Lifetime with Decaborane Between Cleanings

The impact of vapor flow rate on source lifetime (maintenance interval) was studied in a quantitative manner. The electron impact ion source was run continuously with decaborane feed material under controlled conditions at a given vapor flow, until it was determined that the buildup of material was causing a significant decrease in decaborane beam current. Five different flow rates were tested, ranging from about 0.40 sccm to 1.2 sccm. This resulted in mass-analyzed decaborane beam currents (B $_{10}\text{H}_{\star}^{-1}$) ranging from about $150~\mu\text{A}$ to $700~\mu\text{A}$. It is noted that typical feed gas flows in ion sources used in ion implantation range from 1 to about 3 sccm, so this test range is considered a "low" flow regime.

The results of these lifetime tests are summarized in FIG. 5. It suggests a simple model, a hyperbolic function. This is not unexpected; if one assumes zero vapor flow, then the source lifetime would, in principle, diverge; and if one assumes very high vapor flow, then source lifetime would decrease asymptotically to zero. Thus, the model can be expressed by:

(3) (flow rate) × (flow duration) = constant.

Equation (3) simply states that lifetime (i.e., flow duration) is inversely proportional to flow rate; the constant is the amount of deposited material. If equation (3) is accurate, then the fraction of deposited material is independent of the rate of flow of material, which is consistent with our model describing a fixed cross section for dissociation and subsequent deposition. These data show that, using the electron impact ion source with about 0.5 secm decaborane vapor flow, dedicated decaborane operation can be sustained for more than 100 hours. While this is acceptable in many cases, in commercial semiconductor fabrication facilities, source lifetimes of well over 200 hours are desired. When the ion source is used in conjunction with the novel in-situ cleaning procedure of the present invention, greatly extended source lifetimes are achieved.

Advantages of Certain Features of in-situ Ion Source Chemical Cleaning

There are several very important advantages to using a supplemental ion source to produce the reactive gas for in situ cleaning of the ion source. Such plasma sources have been developed for effluent removal applications from process exhaust systems (such as the Litmus 1501 offered by Advanced Energy, Inc.), and for cleaning large CVD process chambers (such as the MKS Astron reactive gas generator), but to the inventors' knowledge it has not been previously recognized that a remote reactive gas generator could be usefully applied to cleaning the ionization chamber of an ion source and the extraction electrode used to generate an ion beam. Remote reactive gas generators such as the MKS Astron have been used to clean process chambers (i.e., relatively large vacuum chambers wherein semiconductor wafers are processed), an application which uses high flows of feed gas (several Standard Liters per Minute (SLM)), and high RF power applied to the plasma source (about 6 kW). The system of the present invention can employ a much more modest feed gas rate, e.g. less than about 0.5 SLM of NP3, and much less RF power (less than about 2.5 kW), for the very small volume of the ionization chamber of the ion source being cleaned (the volume of the ionization chamber for an implanter of semiconductor wafers is typically less than about 100 ml, e.g. only about 75 ml, with a surface area of less than about 200 cm2, e.g. about 100 cm2). The reactive gas flow into the ionization chamber is less than about 2 Standard Liters Per Minute.

One might think it strange to use an external ion source to generate plasma by-products to introduce into the main ion source of the system; why not just introduce the $(e.g., NF_3)$ gas directly into the main ion source to create the plasma by-products within that source directly? The reasons seem not obvious. In order to achieve etch rates which far exceed deposition rates from the feed gas during a small fraction of the uptime (productive period) of the ion implantation system, it is found that the reactive gas must be produced and introduced at relatively very high flows into the small ionization chamber, e.g., flows on the order of 10^2 to 10^3 sccm, compared to typical feed flow rates for the main ion source for ion implantation in the range of 1-3 sccm; such high flows would raise the pressure within the ionization chamber of the main ion source far beyond that for which it is designed to operate for ion implantation. Furthermore, sustaining a high-density NF_3 plasma within the main ion source would etch away sensitive

components, such as hot tungsten filaments. This is because halogen gases etch refractory metals at a high rate which increases exponentially with temperature. (For example, Rosner et al. propose a model for F etching of a tungsten substrate:

(4) Rate (microns/min) = $2.92 \times 10^{-14} \, T^M N_F \, e^{39900T}$,

Where N_F is the concentration of fluorine in atoms per cm³, and T is the substrate temperature in degrees Kelvin.)

Since virtually all ion sources for ion implantation incorporate hot filaments, and since in many cases the ion source chambers are also made of refractory metals such as Mo and W, or graphite (which is aggressively attacked by F), these ion sources would quickly fail under high temperature operating conditions, making the etch cleaning process unusable.

In the presently preferred embodiment, atomic fluorine is caused to enter the cold ionization chamber of the de-energized main ion source at a flow rate of 100 seem or more, and the total gas flow into the ionization chamber is 500 seem or more. Under these conditions, the gas pressure within the ionization chamber is about 0.5 Torr, while the pressure within the vacuum source housing of the implanter is a few tens of milliTorr or more. In a preferred mode of operation, preceding the cleaning phase, an isolation valve is closed between the vacuum housing of the ion source and the implanter vacuum system, and the turbo-molecular pump of the ion source is isolated. The housing of the ion source is then pumped with high-capacity roughing pumps of the vacuum system (i.e., the pumps which normally back the turbo-molecular pumps and evacuate the vacuum system system down to a "rough" vacuum).

A different embodiment of a related etch clean process, shown in FIG. 5, utilizes a "dry etch" gas such as CIF₃. As has previously been observed, the CIF₃ molecule breaks up on contact with deposited surfaces to be cleaned; thus, atomic fluorine and chlorine are released without requiring the generation of a plasma. While handling of CIF₃ gas requires special equipment due to its highly reactive nature, it in principle can simplify the chemical cleaning process to the extent of not requiring an ancillary reactive gas plasma source. Since toxic gases are routinely fed into an ion source for an ion implanter, much of the equipment is already constructed to be "toxic-gas-friendly", and a

separate gas distribution system incorporating CIF₃ can be added in a straightforward manner.

Advantages of the *in-situ* chemical cleaning of the ion source for an ion implanter include: a) extending source life to hundreds, or possibly thousands, of hours before service is required; b) reducing or eliminating cross-contamination brought about by a species change, for example, when switching from octadecaborane ion implantation to arsenic or phosphorus ion implantation, and from arsenic or phosphorus ion implantation to octadecaborane ion implantation; and c) sustaining peak ion source performance during the service life of the ion source.

For example, performing a 10 minute chemical cleaning protocol every eight hours (i.e., once every shift change of operating personnel) and between each species change would have a minimal impact on the uptime of the implanter, and would be acceptable to a modern semiconductor fabrication facility.

Endpoint Detection

It is realized to be beneficial to provide endpoint detection during the cleaning process, so that quantitative information on the efficacy and required duration of the cleaning process may be generated, and the reproducibility of the chemical cleaning process may be assured. FIG. 3 shows a differentially-pumped quadrupole mass analyzer sampling the cleaning process. By monitoring the concentrations of cleaning gas products such as F, Cl, BF₃, PF₃, AsF₃, AlF₃, WF₆, for example, the cleaning process may be tuned and verified. Alternatively, optical means of monitoring the process may be utilized. An FTIR optical spectrometer can monitor the gases resident in the vacuum housing of the ion source of the implanter, through a viewport. This non-invasive (exsitu) means to identify chemical species may be preferable to in-situ monitoring devices in certain cases. Alternatively, see FIG. 4, an extractive FTIR spectrometer may be coupled to the source vacuum housing for endpoint monitoring. A novel means to accomplish endpoint detection consists of monitoring the temperature of the ionization chamber during cleaning. Since the chemical reaction is exothermic, energy is released during the reaction, elevating the chamber temperature. This effect can in principle be used to establish when the reaction rate is diminished.

Novel Ion Extraction Electrode

Borohydrides such as decaborane and octadecaborane are thermally sensitive materials. They vaporize and condense at temperatures between 20C and 100C. It is therefore important to maintain all surfaces with which these materials come into contact at a temperature higher than the vaporizer temperature (but below the temperature at which they dissociate), to prevent condensation. We have found that contamination of the extraction electrode is a problem when using such a borohydride. Both direct ion beam strike and condensed feed vapor or products of its molecular disassociation can degrade operation of the ion beam formation optics, since these boron-containing layers appear to be electrically insulating. Once electrically insulating layers are deposited, they acquire an electrical charge ("charge up") and create vacuum discharges, or "glitches", upon electrical breakdown. Such a discharge creates instabilities in the ion beam current and can contribute to the creation of particles that may reach a process chamber to which the ion beam is directed. An ion implanter which has an ion beam-producing system that experiences many glitches per hour is not considered production-worthy in modern semiconductor fabrication facilities. Furthermore, even in absence of such discharges, as insulating coatings become thicker, the electric charge on electrode surfaces create unwanted stray electric fields which can result in beam steering effects, creating beam loss and may adversely affect ion beam quality.

Discovery of new information has led to a robust solution to this problem. Most implanter ion extraction electrodes are made of graphite. Graphite has been seen to have many advantages in this application, including low materials cost, ease of machining, high electrical conductivity, low coefficient of thermal expansion, and good mechanical stability at high temperatures. However, using a graphite extraction electrode, instabilities were observed after producing an ion beam of borohydrides. It was suspected that the surfaces of the electrode had become insulating. Samples of the electrode deposits were collected and a chemical analysis performed by x-ray fluorescence spectroscopy. The study revealed a chemical stoichiometry consistent with a boron-carbon compound of the form B₂C, which was found to be insulating. In addition, it appeared that metal surfaces in the vicinity of the ion source, including the front plate (i.e., the ion extraction aperture plate) of the ion source also had deposited

insulating coatings after long use. It was conceived to fabricate the electrode of aluminum, and provide radiant heaters to keep the electrode plates, i.e., the suppression and ground electrodes, at a well-controlled, elevated temperature (see Figs. 9, 11) sufficiently high to prevent condensation of decaborane and octadecaborane. In addition, the suppression electrode, which faces the ion source, was fabricated of a single machined piece of aluminum, with a smooth, featureless aspect and all fasteners were located at the backside of the plates. This configuration dramatically reduced the number and severity of discharge points in the event that insulating coatings were formed, employing the principle that the electric field stress at a "point", or sharp feature, is many times greater than at a smooth surface.

The extraction electrode, thus produced, demonstrated excellent performance, and operated reliably for more than 100 hours (at least ten times as long as the graphite electrode) with very low glitch frequency. This great improvement is attributed to: i) Al construction (i.e., metal versus graphite), ii) Active heating and temperature control of the electrode plates, and iii) smooth electrode surfaces. It was found that operating the electrode plates at 200C gave good results when running decaborane, significantly reducing the amount of deposited material. In general, the temperature of the extraction electrode should be kept below the dissociation temperature of the feed material. In the case of decaborane the temperature should be kept below 350C, preferably in the range 150C to 250C. For octadecaborane operation, the temperature should not exceed 160C, since chemical changes occur in octadecaborane above this temperature; when running octadecaborane, an extraction electrode temperature between 120C and 150C yields good results.

The radiative design shown in FIG. 11 demonstrated very good temperature uniformity. Incorporating resistive heaters, particularly using an aluminum electrode as illustrated in FIG. 12, can also yield good uniformity and results in a more compact design requiring less maintenance.

Por constructing a heated extraction electrode, other metals would also work, for example molybdenum. Molybdenum has the advantage of being refractory, so it can withstand very high temperatures. It also has good thermal conductivity. Aluminum, on the other hand, is a column III element like In and B of the periodic table, and therefore

offers the advantage of being only a mild contaminant in silicon (it is a P-type dopant in silicon), while transition metals such as molybdenum are very detrimental to carrier lifetimes in integrated circuits. Aluminum is also not readily attacked by halogens, whereas transition metals such as molybdenum are susceptible to attack, particularly at elevated temperatures. The primary disadvantage of aluminum, however, is that it is not a high temperature material, and should be used below about 400 C.

For these reasons, depending upon the particular use, the heated electrode is constructed of a selected heat-resistant material, aluminum or an aluminum containing complex often being preferred when used in association with in situ etch cleaning.

BRIEF DESCRIPTION OF THE DRAWINGS

- FIG. 1: Ion beam generation system incorporating reactive gas cleaning.
- FIG. 2: Second embodiment of ion beam generation system incorporating reactive gas cleaning.
- FIG. 3: Ion beam generation system similar to FIG. 1 but incorporating a vaporizer and certain gas distribution elements.
- FIG. 4: Ion beam generation system similar to FIG. 2 but incorporating a vaporizer and certain gas distribution elements.
- FIG. 5: Ion generation system incorporating reactive gas cleaning by the introduction of CIF₃.
- FIG. 6: Gas box for an ion implanter which includes a reactive gas plasma source, feed vapor source, ion source electronics, and facilities for the plasma source.
 - FIG. 6A: View similar to Fig. 6, showing a vapor flow control system .
 - FIG. 6B: Valve schematic for an ion beam generating system.
 - FIG. 7: Electron-impact ion source.
 - FIG. 7A: Magnified view of a portion of FIG. 7, showing shielding of elements.
 - FIG. 7B: Control diagram for an embodiment.
 - FIG. 8: Ion extraction electrode.
 - FIG. 9: Ion extraction electrode optics.
 - FIG. 9a: B₁₈H_x⁺ beam profiles.
 - FIG. 10: Extraction electrode and manipulator.

FIG. 11: Electrode head-exploded view.

FIG. 12: Second embodiment of electrode head.

FIG. 13: B₁₀H_x⁺ beam current versus decaborane flow rate.

FIG. 14: Lifetime versus decaborane vapor flow rate.

FIG. 15: Etch rate of Si coupon.

FIG. 16: Ion implanter.

DETAILED DESCRIPTION

Novel Ion Beam-Generating System

FIG. 1 shows an ion beam-generating system. As shown in this example, it is adapted to produce an ion beam for transport to an ion implantation chamber for implant into semiconductor wafers or flat-panel displays. Shown are ion source 400, extraction electrode 405, vacuum housing 410, voltage isolation bushing 415 of electrically insulative material, vacuum pumping system 420, vacuum housing isolation valve 425, reactive gas inlet 430, feed gas and vapor inlet 441, vapor source 445, feed gas source 450, reactive gas source 455, ion source high voltage power supply 460, and resultant ion beam 475. An ion beam transport housing is indicated at 411. The ion source 400 is constructed to provide cluster ions and molecular ions, for example the borohydride ions B₁₀H_x⁺, B₁₀H_x⁻, B₁₈H_x⁺, and B₁₈H_x or, or in addition, more conventional ion beams such as P+, As+, B+, In+, Sb+, Si+, and Ge+. Ion source 400 may be a Bernas-style arcdischarge ion source, which is most commonly used for ion implantation, or a "bucket"type water-cooled ion source which uses an immersed RF (radio frequency) antenna forming an RF field to create ions, a microwave ion source, or an electron-impact ionization source, for example. The gas and vapor inlet 441 for gaseous state feed material to be ionized is connected to a suitable vapor source 445, which may be in close proximity to gas and vapor inlet 441 or may be located in a more remote location, such as in a gas distribution box located elsewhere within a terminal enclosure. A terminal enclosure is a metal box, not shown, which encloses the ion beam generating system. It contains required facilities for the ion source such as pumping systems, power distribution, gas distribution, and controls. When mass analysis is employed for selection

of an ion species in the beam, the mass analyzing system may also be located in the remnial enclosure.

In order to extract ions of a well-defined energy, the ion source 400 is held at a high positive voltage (in the more common case where a positively-charged ion beam is generated), with respect to the extraction electrode 405 and vacuum housing 410, by high voltage power supply 460. The extraction electrode 405 is disposed close to and aligned with the extraction aperture 504 of the ionization chamber. It consists of at least two aperture-containing electrode plates, a so-called suppression electrode 406 closest to ionization chamber 500, and a "ground" electrode 407. The suppression electrode 406 is biased negative with respect to ground electrode 407 to reject or suppress unwanted electrons which are attracted to the positively-biased ion source 400 when generating positively-charged ion beams. The ground electrode 407, vacuum housing 410, and terminal enclosure (not shown) are all at the so-called terminal potential, which is at earth ground unless it is desirable to float the entire terminal above ground, as is the case for certain implantation systems, for example for medium-current ion implanters. The extraction electrode 405 may be of the novel temperature-controlled metallic design, described below.

(If a negatively charged ion beam is generated the ion source is held at an elevated negative voltage with other suitable changes, the terminal enclosure typically remaining at ground.)

FIG. 1 shows the reactive gas source 455 at terminal potential, with reactive gas inlet 430 incorporating a high voltage break 431, which can be fabricated of an insulating ceramic such as $\mathrm{Al}_2\mathrm{O}_3$, for example. Since ion sources for ion implantation can in general be biased up to a maximum voltage of about 90 kV, this high voltage break 431 must stand off 90 kV for that application. As will be described below, the cleaning system is used only with the ionizing source and high voltages off (de-energized), so that there is only high voltage across break 431 when the vacuum housing 410 is under high vacuum, which makes high voltage standoff clearance requirements easier to meet. A dedicated endpoint detector 470, in communication with the vacuum housing 410, is used to monitor the reactive gas products during chemical cleaning.

For ion sources suitable for use with ion implantation systems, e.g. for doping semiconductor wafers, the ionization chamber is small, having a volume less than about 100 ml, has an internal surface area of less than about 200cm², and is constructed to receive a flow of the reactive gas, e.g. atomic fluorine or a reactive fluorine-containing compound at a flow rate of less than about 200 Standard Liters Per Minute.

It is seen that the system of FIG. 1 enables in situ cleaning, i.e. without the ion source being removed from its operating position in the vacuum housing, and with little interruption of service.

FIG. 2 illustrates another embodiment. The principal difference in FIG. 2 over FIG. 1 is that the reactive gas source 455 and reactive gas inlet 430 are at ion source potential. The benefits of this approach are twofold: it is a more compact arrangement, and it allows the reactive gas source 455 and its associated gas supplies to be contained in the gas box which, at ion source potential supplies gas and power to the ion source 400, as is typical in commercial ion implantation systems.

Chemical Cleaning System

The embodiment of FIG. 3, having many features similar to FIG. 1, is constructed to generate, selectively, both cluster ions and monomer ions. It has a dedicated gas inlet 435 for feed material in normally gaseous state and is in communication, through valve 443, with a vapor source 445 for producing borohydride and other vaporized feed materials. For conducting in-situ chemical cleaning of the ion source and electrode, a remote plasma source 455 disassociates gas supplied by a cleaning gas supply 465, for example NF₃, into decomposition products such as F, F₂, and N-containing compounds. When cleaning is desired, after de-energizing the ion source, the decomposition products are fed into the ionization chamber from the outlet 456 of the remote plasma source 455 by dedicated reactive gas inlet 430. The remote plasma source 455 is mounted on the terminal potential side of voltage isolation bushing 415. Since the ion source 400 runs at high voltage, a high voltage break 431 in vacuum provides voltage isolation.

To initiate a cleaning cycle, the ion source is shut down and vacuum housing isolation valve 425 is closed; the high vacuum pump 421 of the vacuum pumping system 420 is isolated and the vacuum housing 410 is put into a rough vacuum state of <1 Torr

by the introduction of dry N_2 gas while the housing is actively pumped by backing pump 422. Once under rough vacuum, argon gas (from Ar gas source 466) is introduced into the plasma source 455 and the plasma source is energized by on-board circuitry which couples radio-frequency (RF) power into the plasma source 455. Once a plasma discharge is initiated, Ar flow is reduced and the F-containing cleaning gas feed 465, e.g. NF₂, is introduced into plasma source 455. Reactive F gas, in neutral form, and other byproducts of disassociated cleaning gas feed 465, are introduced through reactive gas inlet 430 into the de-energized ionization chamber 500 of ion source 400. The flow rates of Ar and NF₃ (for example) are high, between 0.1 SLM (Standard Liters per Minute) and a few SLM. Thus, up to about 1 SLM of reactive F as a dissociation product can be introduced into the ion source 400 in this way. Because of the small volume and surface area of ionization chamber 500, this results in very high etch rates for deposited materials. The ionization chamber 500 has a front plate facing the extraction electrode, containing the extraction aperture 504 of cross sectional area between about 0.2 cm2 and 2 cm2, through which, during energized operation, ions are extracted by extraction electrode 405. During cleaning, the reactive gas load is drawn from ionization chamber 500 through the aperture 504 by vacuum of housing 410; from housing 410 the gas load is pumped by roughing pump 422. Since the extraction electrode 405 is near and faces aperture 504 of ionization chamber 500, the electrode surfaces intercept a considerable volume of the reactive gas flow. This results in an electrode cleaning action, removing deposits from the electrode surfaces, especially from the front surface of suppression electrode 406, which is in position to have received the largest deposits. Thus, it is beneficial to fabricate extraction electrode and its mounting of F-resistant materials, such as Al and Al2O3.

The embodiment of FIG. 3 also has an endpoint detector consisting of a differentially-pumped, Residual Gas Analyzer (RGA), constructed for corrosive service. Analyzer RGA is in communication with vacuum housing 410. It is to be used as a detector for the end point of the cleaning action by monitoring partial pressures of F-containing reaction products (for example, BF3 gas resulting from B combining with F). Other types of endpoint detectors can be used, the RGA being shown to illustrate one particular embodiment. When the boron-containing partial pressures decline at RGA, the

. PCT/US2004/041525

cleaning process is largely completed. Once the cleaning process is ended, the plasma source 455 is turned off and is briefly purged with Ar gas (which also purges the ionization chamber 500, the housing 410 and elements contained therein). The roughing pump 422 is then isolated from direct communication with vacuum housing 410, the high vacuum pump 421 isolation valve is opened, and vacuum housing 410 is restored to high vacuum (about 1×10^{6} Torr or below). Then, vacuum housing isolation valve 425 is opened. The system is now ready to resume ion beam generation. The ion source voltage supply 460 can be energized and ion source 400 operated normally.

An advantage of the embodiment of FIG. 3 is that the service facilities needed to support the remote plasma source 455, such as cooling water circulation and electrical power, can be at the terminal potential of an ion implanter (see 208 in FIG. 16). This enables sharing facilities denoted at S such as cooling water and electrical power, with the mass-analyzer magnet 230 of the implanter. During cleaning mode, when plasma source 455 is energized, the analyzer 230 is de-energized and therefore does not need water or power, and vise versa, during ion beam production mode. This "sharing" can be accomplished by suitable control arrangements represented diagrammatically at S', which direct service facilities such as cooling water circulation and power supply connection alternatively to the analyzer magnet 230, dashed arrow S, or to the remote plasma source 455, solid arrow S, depending upon the mode of operation being employed.

FIG. 4 shows an implementation similar to FIG. 2 for conducting *in-situ* chemical cleaning of an source 400 and extraction electrode 405. Three inlet passages are integrated into ion source 400, respectively for reactive gas 430 from plasma source 455, feed gas 435 from one of a number of storage volumes 450 selected, and feed vapor 440 from vaporizer 445. Unlike FIG. 3, the embodiment of FIG. 4 has the plasma-based reactive gas source 455 at the high voltage of ion source 400. This enables the remote plasma source 455 to share control points of the ion source 400, and also enables the cleaning feed gas 465 and argon purge gas from storage 466 to be supplied from the ion source gas distribution box, which is at source potential, see also FIGs. 6 and 6A. Also shown is a different type of endpoint detector, namely a Fourier Transform Infrared (FTIR) optical spectrometer. This detector can function ex-situ (outside of the vacuum

housing), through a quartz window. Instead, as shown in FIG. 4, an extractive type of FTIR spectrometer may be used, which directly samples the gas in the vacuum housing 410 during cleaning. Also a temperature sensor TD may sense the temperature of the deenergized ionization chamber by sensing a thermally isolated, representative region of the surface of the chamber. The sensor TD can monitor heat produced by the exothermic reaction of F with the contaminating deposit, to serve as an end-point detection.

FIG. 5 shows an ion beam-generating system similar to that of FIG. 4, but incorporating a fundamentally different type of reactive gas source 455. In this case, reactive CIF₃ gas contained in a gas cylinder is fed directly into ion source 400 without use of a remote plasma source. This potentially reduces equipment cost and footprint since power and controls for a remote plasma source are not required. However, since CIF₃ is pyrophoric, it is dangerous and requires special gas handling, whereas NF₃ (for example) is primarily an asphyxiant, and is less toxic than many semiconductor gases, such as BF₃, PH₃, or AsH₃, and therefore safer.

FIG. 6 shows plasma source 455, vapor source 445, source electronics, and service facilities S for the plasma source contained within a gas box B meant for retrofit into an existing ion implanter installation.

The embodiment of FIG. 6a differs from the embodiment of FIG. 6 above, by incorporating a preferred vaporizer and flow control system described below. FIG. 6B is a valve schematic diagram for the ion source and self-cleaning system of FIG. 4.

Preferred Ion Source and Vaporizer

FIG. 7 is a diagram of a preferred ion source 10 and its various components, and see FIG. 7A. The details of its construction, as well as its preferred modes of operation, are similar to that disclosed by Horsky et al., International Application No. PCT/US03/20197, filed June 26, 2003: "An ion implantation device and a method of semiconductor manufacturing by the implantation of boron hydride cluster ions", and by Horsky, U.S. Patent Application No. 10/183,768, "Electron impact ion source", filed June 26, 2002, both herein incorporated by reference. The ion source 10 is one embodiment of a novel electron impact ionization system. Figure 7 is a cross-sectional schematic diagram of the source construction which serves to clarify the functionality of the

components which make up the ion source 10. The ion source 10 is made to interface to an evacuated vacuum chamber of an ion implanter by way of a mounting flange 36. Thus, the portion of the ion source 10 to the right of flange 36, shown in FIG. 7, is at high vacuum (pressure <1×10⁻⁴ Torr). Gaseous material is introduced into ionization chamber 44 in which the gas molecules are ionized by electron impact from electron beam 70, which enters the ionization chamber 44 through electron entrance aperture 71 such that electron beam 70 is aligned with (i.e. extends adjacent, parallel to) ion extraction aperture 81. Thus, ions are created adjacent to the ion extraction aperture 81, which appears as a slot in the ion extraction aperture plate 80. The ions are then extracted and formed into an energetic ion beam 475 by an extraction electrode 220 (FIGs. 8 and 9) located in front of the ion extraction aperture plate 80. Referring to FIG. 7, gases such as argon, phosphine, or arsine, for example, may be fed into the ionization chamber 44 via a gas conduit 33. Solid feed materials 29 such as decaborane or octadecaborane can be vaporized in vaporizer 28, and the vapor fed into the ionization chamber 44 through vapor conduit 32 within the source block 35. Typically, ionization chamber 44, ion extraction aperture plate 80, source block 35 (including vapor conduit 32), and vaporizer housing 30 are all fabricated of aluminum. Solid feed material 29 is held at a uniform temperature by closed-loop temperature control of the vaporizer housing 30. Sublimated vapor 50 which accumulates in a ballast volume 31 feeds through conduit 39 and through throttling valve 100 and shutoff valve 110. The nominal pressure of vapor 50 between throttling valve 100 and shutoff valve 110 is monitored by heated pressure gauge 60, preferably a capacitance manometer. Since the vapor 50 feeds into the ionization chamber 44 through the vapor conduit 32, located in the source block 35, and gases feed in through gas conduit 33, both gaseous and vaporized materials may be ionized by this ion source, which is capable of creating ion beam 475 consisting of either molecular ions (such as B₁₈H_x⁺)or monomer ions (such as As⁺), as needed.

Vapor Flow Control Into the Ion Generating System

The flow of vapor to ionization chamber of FIG. 7, and see FIG. 7B, is determined by the vapor pressure in the region just before vapor feed passage 32, *i.e.*, within shutoff valve 110 in FIG. 7. This is measured by pressure gauge 60, e.g. a

capacitance monometer, located between throttling valve 100 and shut-off valve 110. In general, the flow rate is proportional to the vapor pressure. This allows the pressure signal to represent flow, and to be used as a set point to select flow. To generate a desired vapor flow into the ion source, vaporizer housing 30 is brought to a temperature such that when throttling valve 100 is in its fully open position, the desired flow rate is exceeded. Then the throttling valve 100 is adjusted to reach the desired pressure output.

To establish a stable flow over time, separate closed loop control of the vaporizer temperature and vapor pressure is implemented using dual PID controllers, such as the Omron E5CK control loop digital controller. The control (feedback) variables are thermocouple output for temperature, and gauge output for pressure. The diagram of FIG. 7B shows a digital vapor feed controller 220 for performing these closed loop control functions.

In FIG. 7B gauge output 250 from pressure gauge 60 is applied to throttle valve position control 245 which applies throttle valve position control signal 247 to throttle valve 100. Thermocouple output 225 from vaporizer 28 is applied to vaporizer heater control 215 which controls heater power 248 applied to the vaporizer 28.

A second, slow level of control is implemented by digital feed controller 220, accommodating the rate at which solid feed material vaporizes being a function of its open surface area, particularly the available surface area at the solid-vacuum interface. As feed material within the vaporizer is consumed over time, this available surface area steadily decreases until the evolution rate of vapors cannot support the desired vapor flow rate, resulting in a decrease in the vapor pressure upstream of the throttle valve 100. This is known as "evolution rate limited" operation. So, with a fresh charge of feed material in the vaporizer, a vaporizer temperature of, say, 25C might support the required vapor flow at a nominal throttle valve position at the low end of its dynamic range (i.e., the throttling valve only partially open). Over time (for example, after 20% of the feed material is consumed), the valve position would open further and further to maintain the desired flow. When the throttle valve is near the high conductance limit of its dynamic range (i.e., mostly open), this valve position is sensed by the controller 220, which sends a new, higher heater set point temperature to the vaporizer heater control 215. The increment is selected to restore, once the vaporizer temperature settles to its new value,

the nominal throttle valve operating point near the low end of its dynamic range. Thus, the ability of the digital controller 220 to accommodate both short-timescale changes in set point vapor pressure and long-timescale changes in vaporizer temperature makes the control of vapor flow over the lifetime of the feed material charge very robust. Such control prevents over-feeding of vapor to the ionization chamber. This has the effect of limiting the amount of unwanted deposits on surfaces of the ion generating system, thus extending the ion source life between cleanings.

FIG. 8 shows a top view (looking down) of an ion extraction electrode 220 facing the novel ion source 10. The ion source 10 is held at a positive potential V_A with respect to the ion extraction electrode 220, which is at local ground potential, *i.e.*, at the potential of the vacuum housing. The ion extraction electrode 220 is a simple diode; electrode plate 302 is the "ground" electrode and plate 300 the "suppression" electrode, typically held a few thousand volts below ground potential by suppression power supply V_S . The ionization chamber 44 and ion extraction aperture plate 80 of ion source 10 are shown facing extraction electrode 220. The three plates 80, 300, 302 contain rectangular slots or apertures through which ions 90 are extracted; FIG. 8 illustrates the slot profiles in the "short", or dispersive, direction.

Novel Heated Electrode

During the decaborane lifetime tests shown in FIG. 14, a novel heated aluminum electrode was used. FIG. 9 shows a top view of the basic optical design of the extraction system, in the dispersive plane of the one-dimensional "slot" aperture lenses. In the implanter used, the ionization chamber 490 of the ion source was held at the desired ion beam energy by positive high voltage power supply V_A . Fig. 8. For example, if a 20 keV ion beam is desired, then $V_A = 20$ keV. Ion extraction aperture plate 500 is electrically isolated from ionization chamber 490 such that it can be biased by bipolar power supply V_B from .750V—750V. The isolation is accomplished by a thermally conductive, electrically insulating polymeric gasket which is sandwiched between the ion extraction aperture plate 500 and ionization chamber 490. The parts of the ion source body that are exposed to vapor (source block 35, ionization chamber 44, and extraction aperture plate 80 in FIG. 7) are maintained in good thermal contact with each other to maintain

controlled temperature surfaces during source operation. Ions produced in ionization chamber 490 are extracted through the aperture in ion extraction aperture plate 500 by extraction electrode 540 consisting of suppression electrode 510 and ground electrode 520. The ions propagate as a focused ion beam along the beam axis 530. Suppression electrode 510, biased to a few thousand volts negative by power supply Vs, serves to suppress secondary electrons which are generated upstream from the suppression electrode due to beam strike, preventing these energetic electrons from backstreaming into the positively-biased ion source. The ionization chamber 490, ion extraction aperture plate 500, suppression electrode 510, and ground electrode 520 are all fabricated of aluminum, and have smooth, carefully polished surfaces to minimize local electric fields.

An important effect of biasing ion extraction aperture plate 500 is to change the focal length of the ion optical system of FIG. 9. A negative bias increases the focal length, while a positive bias decreases the focal length. For large biases, the effect can be substantial. For diagnostic purposes, a scanning-wire profilometer was installed, located at the entrance to the analyzer magnet, just downstream of the source housing isolation valve (210 in FIG. 16). This scanner recorded the beam current distribution in the dispersive plane, useful to determine how well the ion beam is being focused in the dispersive plane. 20 keV octadecaborane beam profiles are shown in FIG. 9a for three different biasing conditions: -483V, 0, and +300V. The zero volt condition is substantially overfocused, the positive voltage condition more overfocused, and the negative voltage condition properly focused. The electrode position was held constant during the three measurements. As expected, the proper focusing condition yielded the highest ion beam currents.

The ability to change the optical focal length, and thus tune the optical system to obtain the highest ion beam current, enables introduction of the least amount of feed material to the vaporizer. Again, this has the beneficial effect of limiting the amount of unwanted deposits on surfaces of the ion generating system, extending the ion source life between cleanings.

Besides the biasing of the extraction aperture plate for focusing the system just described, the invention provides means for moving the extraction electrode optic

element relative to other components of the system. FIG. 10 shows the novel electrode 600 mounted on a three-axis manipulator 610 which allows for motion (with respect to the ion source) in X, Z and Θ , as defined by coordinate system 620. Actuator 613 controls X-motion, actuator 612 controls Z-motion, and actuator 611 controls Θ -motion. The manipulator 610 mounts to the side of the implanter vacuum housing via mounting flange 615.

FIG. 11 shows a partial exploded view of the radiatively-heated version of the novel electrode head. Shown are suppression electrode 700, ground electrode 710, heater plate 720, and radiant heater wire 730. The suppression and ground electrodes are fabricated of aluminum, the heater plate of stainless steel, and the heater wire 730 of nichrome. When the electrode was operated at 200C, power consumption was about 60W to maintain the temperature. The heater power is controlled with a closed-loop PID controller, the Omron E5CK, based on readback of a thermocouple.

FIG. 12 shows a partial exploded view of a resistively-heated version of the novel electrode head. Shown are suppression electrode 800, ground electrode 810, and resistive heaters 820. The four resistive heaters 820 fit into sleeves 830, two into each electrode plate. The sleeves 830 are a split design such that the heater press-fits into the sleeve, achieving intimate contact. Intimate contact between heater and electrode is important to insure proper heating of the electrode, and to prevent premature burnout of the heaters. Again, the Omron E5CK or equivalent can control the electrode temperature based on readback of a thermocouple.

As described above, use of these heating arrangements for the extraction electrode maintain a well-controlled, elevated temperature sufficiently high to prevent condensation of decaborane and octadecaborane such as produced by the relatively cool-operating ion source of FIGs. 7 and 7A. The extraction electrode made of fluorine-resistant materials, e.g. aluminum, enables periodic in situ cleaning of the electrode to remove any deposits by fluorine vapors drawn through the extraction aperture.

A different situation is encountered with plasma ion sources that inherently run so hot that the heat may harm the extraction electrode assembly if made of low temperature material. Referring to Fig. 9, shown in dotted lines are circular cooling coils, 512 and 522 secured in heat transfer relationship to the backs of aluminum electrode members

510 and 522, respectively. Circulation of cooling fluid through these cooling coils can cool the aluminum electrodes to prevent deformation by heat from hot ion sources. This enables use of fluorine-resistant materials for the extraction electrode, for instance aluminum or a complex containing aluminum, which provide resistance to attack by any fluorine present from feed materials or from reactive cleaning gas.

Source Lifetime Measurements When Running Decaborane

FIG. 14 shows the results of source lifetime testing over a broad range of decaborane flows. The fit to these data is from Equation (3). No failures of the ion source were recorded during these tests; rather, the individual tests were ended when the decaborane ion current dropped to roughly half of its initial level. Upon inspecting the ion source, it was found that a substantial amount of boron-containing material was deposited within the ionization chamber, mostly adhering to the interior walls of the chamber. In some cases, the ion extraction aperture was also partially occluded. The model of Equation (3) seems to fit the data well, and suggests that "lean" operation is the key to prolonged ion source lifetime, between in situ chemical cleaning procedures or disassembly.

Measurements of Etch Rates Within Ionization Chamber During F Cleaning

The system with the ion source 10 of FIG. 7 was used to test the F cleaning process on 1-mm-thick silicon coupons staged inside of the ionization chamber 44, with the following modification: rather than incorporating a dedicated reactive feed conduit, the vapor feed conduit 32 was employed to introduce the reactive gas. Si was used because etching of Si by F is well understood, and pure Si material is available in the form of Si wafers. This test required removing the vaporizer between cleaning cycles. Two coupon locations were tested: one having line-of-sight relationship with the reactive gas inlet (i.e., the vapor feed 32), and one not having line-of-sight. The etch rates are shown in FIG. 15 as a function of NF₂ flow rate. During this process, a flow of 700 sccm of argon was maintained into the remote plasma source while the NF₂ flow rate was varied from 50 sccm to 500 sccm. A line-of-sight geometry shows a factor of about five increase in etch rate, and is therefore a preferred geometry if it can be done uniformly.

To this end, the geometry portrayed in FIG. 3 should provide better etch uniformity of the ion source ionization chamber 44 than the geometry shown in FIG. 4: The test also indicated that location of etch-sensitive components shielded from the gas flow is effective to provide a degree of protection to those components.

To extend the life of components of the self-cleaning ion generating system construction materials are selected that are resistive to the reactive gas, and provision can be made for shielding of sensitive components.

For the interior of the ionization chamber, as indicated above, aluminum is employed where the temperature of the ionizing action permits because aluminum components can withstand the reactive gas fluorine. Where higher temperature ionizing operation is desired, an aluminum-silicon carbide (AlSiC) alloy is a good choice for the surfaces of the ionization chamber or for the extraction electrode. Other materials for surfaces in the ionization chamber are titanium boride (TiB₂), Boron Carbide (B₄C) and silicon carbide (SiC).

For components exposed to the fluorine but not exposed to the ionizing action, for instance electron source components such as electrodes, the components may be fabricated of Hastelloy, fluorine-resistant stainless steels and nickel plated metals, for instance nickel-plated molybdenum.

Both inert gas shields and movable physical barriers can protect components of the system from the reactive gas during cleaning. For example, referring to FiG. 7A, a conduit 113 for inert gas, for instance argon, extends from a gas source, not shown. Its outlet is at a strategic location in the ion source, such that flow of the inert gas, when initiated for the cleaning cycle, floods the component to be protected. In FiG. 7A the outlet 113a of inert gas conduit 113 aims a flooding stream of argon over the active components of electron gun 112, including, the electron-emitting cathode. In FiG. 7A a movable shield member 73 is also shown, which is movable into position for the cleaning cycle. In the example shown, it is movable over the aperture 71A leading to beam dump 72, or to another electron gun when provided on that side of the ionization chamber 44.

The cleaning process described above was conducted to observe its effect on boron deposits within the ionization chamber and on the interior of the ion extraction

aperture of the novel ion source 10 of FIG. 7. The observed etch rates had characteristics similar to the plot of FIG. 15, but were a factor of 3 lower. Thus, for a NF3 flow rate of 500 sccm, the etch rate for decaborane deposits were 7 μ m/min (no line-of-sight), and 36 μ m/min (line-of-sight). The interior of the ion extraction aperture after running 4 hrs of decaborane at 0.8 sccm vapor flow had about 133 μ m thick boron-containing deposit prior to cleaning. Observations were made after a 5 min F clean, and after a 15 min F clean using these flow rates. One side of the aperture plate was in line-of-sight with the vapor feed. It was observed from the cleaning pattern that the vapor feed aperture is centered in the vertical direction! After 15 minutes of cleaning, the plate was almost completely free of deposits. Also, the novel heated aluminum ion extraction electrode of FIG. 10 was removed and inspected after long operation. It was very clean with no observable decaborane deposits. This was undoubtedly due to exposure of the electrode to reactive F (F can flow through the ion source ion extraction aperture located in front of the vapor conduit, to the extraction electrode directly in front of it). Also, elevated temperature of the Al electrode assembly increased the effective etch rate of its deposits.

With respect to the ionization chamber, again, a 15 min etch clean left the chamber nearly free of deposits. A test was conducted in which the system was repeatedly cycled in the following manner: two hours of decaborate operation (>500μA of analyzed beam current), the source was turned off and the filament allowed to cool, followed by a 15 min chemical clean at 500sccm of NF₃ feed gas and 700sccm of Ar, to see if conducting repeated chemical cleaning steps was injurious to the ion source or extraction electrode in any-way. After 21 cycles there was no measurable change in the operating characteristics of the ion source or the electrode. This result demonstrates that this F cleaning process enables very long lifetime in ion source operation of condensable species.

The Ion Generating System Incorporated Into an Exemplary Ion Implanter

FIG. 16 shows the basic elements of a commercial ion implanter, with an embodiment of the novel ion beam generation system incorporating the ion source of FIG. 7 installed. The ion source 10 is inserted into the source vacuum housing 209 of the ion implanter. It is electrically insulated from housing 209 by insulator 211. The ion

extraction electrode 220 extracts and accelerates ions from the ion source 10 to form an ion beam 200. Ion beam 200 propagates entirely in vacuum; from the electrode 220 it enters analyzer housing 290, 300 where it is bent and dispersed by dipole analyzer magnet 230 into separate beamlets which differ by their charge-to-mass ratio. The ion beamlet of interest passes through mass resolving aperture 270 and into a final acceleration (or deceleration) stage 310. The thus-produced, selected and-accelerated ion beam 240 leaves the ion beam forming system 208 and is introduced to the process chamber 330 where it intercepts one or more device wafers 312 on rotating disk 314. The ion source vacuum housing 209 can be isolated from the remainder of the implanter's vacuum system by closing isolation valve 210. For example, isolation valve 210 is closed prior to in situ cleaning of the ion source.

CLAIMS:

 A system for generating an ion beam (475 or 240) comprising an ion source (400 or 10) in combination with an extraction electrode (405 or 220) and a reactive gas cleaning system (455, 430),

the ion source comprising an ionization chamber (500) connected to a high voltage power supply and having an inlet (441,440 or 435) for gaseous or vaporized feed materials, an energizeable ionizing system for ionizing the feed material within the ionization chamber and an extraction aperture (304) that communicates with a vacuum housing (410 or 209), the vacuum housing evacuated by a vacuum pumping system (420).

the extraction electrode (405 or 220) disposed in the vacuum housing outside of the ionization chamber, aligned with the extraction aperture (504) of the ionization chamber and adapted to be maintained at a voltage below that of the ionization chamber to extract ions through the aperture from within the ionization chamber,

and the reactive gas cleaning system (455,430) operable when the ionization chamber and ionizing system are de-energized to provide a flow of reactive gas through the ionization chamber (500) and through the ion extraction aperture (504) to react with and remove deposits on at least some of the surfaces of the ion generating system.

- .2 The system for generating an ion beam of claim 1 constructed for use in implanting ions in semiconductor wafers, the ionization chamber (500) having a volume less than about 100ml and an internal surface area of less than about 200cm².
- 3. The system for generating an ion beam of claim 1 or 2 constructed to produce a flow of the reactive gas into the ionization chamber at a flow rate of less than about 2 Standard Liters Per Minute.
- 4. The system for generating an ion beam of any of the foregoing claims in which the extraction electrode (405 or 220) is constructed to produce a beam of accelerated ions suitable for transport to a point of utilization.

5. The system for generating an ion beam of any of the foregoing claims in which the extraction electrode (405 or 220) is located within a path of reactive gas moving from the extraction aperture (504) to the vacuum pumping system so that the extraction electrode is cleaned by the reactive gas.

- 6. The system for generating an ion beam of any of the foregoing claims in which the extraction electrode (405 or 220) is associated with a heater (720, 730, 820) to maintain the electrode at elevated temperature during extraction by the extraction electrode of ions produced in the ionization chamber, e.g. above the condensation temperature, below the disassociation temperature of solid-derived, thermally sensitive vapors.
- 7. The system of any of the foregoing claims 1-5 in which the extraction electrode (405; 220) is associated with a cooling device (512, 522), e.g. when the electrode is formed of temperature sensitive material and is used with a hot ion source.
- The system for generating an ion beam of any of the foregoing claims in which
 the extraction electrode has a smooth, featureless aspect.
- 9. The system for generating an ion beam of any of the foregoing claims in which the reactive gas cleaning system (455, 430) comprises a plasma chamber, the plasma chamber arranged to receive a feed gas capable of being disassociated by plasma to produce a flow of reactive gas through a chamber outlet (456), and a conduit (430) for transporting the reactive gas to the ionization chamber (500).
- 10. The system for generating an ion beam of claim 9 in which the plasma chamber (500) is constructed and arranged to receive and disassociate a compound capable of being disassociated to atomic fluorine, for instance NF₃, C₃F₈ or CF₄.
- 11. The system for generating an ion beam of claim 9 or 10 in which the reactive gas cleaning system is constructed and arranged to share a service facility (S) associated with the ion source (400 or 10).

12. The system for generating an ion beam of claim 9 or 10 constructed to direct an ion beam through a mass analyzer (230), in which the reactive gas cleaning system is constructed and arranged to share a service facility (S) with the mass analyzer.

- 13. The system for generating an ion beam of any of the foregoing claims 1-8 in which the reactive gas cleaning system comprises a conduit (430A) from a container of pressurized reactive gas, for instance C1F₃.
- · 14. The system for generating an ion beam of any of the foregoing claims in combination with an end-point detection system (470, RGA, FIIR, TD) adapted to at least assist in detecting substantial completion of reaction of the reactive gas with contamination on a surface of the system for generating an ion beam.
 - 15. The system for generating an ion beam of claim 14 in which the end point detection system comprises an analysis system (for instance, RGA or FIIR) for the chemical makeup of gas that has been exposed to said surface during operation of the reactive gas cleaning system.
 - 16. The system for generating an ion beam of claim 14 or 15 including a temperature detector (TD) arranged to detect substantial termination of an exothermic reaction of the reactive gas with contamination on a surface of the system.
 - 17. The system for generating an ion beam of any of the foregoing claims in which the energizeable ionizing system includes a component within or in communication with the ionization chamber (500) that is susceptible to harm by the reactive gas and means are provided to shield the component from reactive gas flowing through the system.
 - 18. The system for generating an ion beam of claim 17 in which the means comprises an arrangement for producing a flow of inert gas, such as argon, past the component.

19. The system for generating an ion beam of claim 17 or 18 in which the means comprises a shield member that is impermeable to the reactive gas.

- 20. The system for generating an ion beam of any of the foregoing claims constructed to operate with reactive halogen gas as the reactive gas and the extraction electrode (405 or 220) and associated parts comprise aluminum (A1) or alumina (Al₂O₃).
- 21. The system for generating an ion beam of any of the foregoing claims in which the ion source (400 or10) is constructed to produce ions within the ionization chamber (500) via an arc-discharge, an RF field, a microwave field or an electron beam.
- 22. The system for generating an ion beam of any of the foregoing claims associated with a vaporizer (28, 445) of condensable solid feed material for producing feed vapor to the ionization chamber.
- 23. The system for generating an ion beam of any of the foregoing claims in which the ion source is constructed to vaporize feed material capable of producing cluster or molecular ions, and the ionization system is constructed to ionize the material to form cluster or molecular ions for implantation.
- 24. The system for generating an ion beam of any of the foregoing claims in which the vacuum housing (410 or 209) is associated with a pumping system (420) comprising a high vacuum pump (421) capable of producing high vacuum and a backing pump (422) capable of producing rough vacuum, the high vacuum pump (421) operable during operation of the ion source (400 or 10), and being capable of being isolated from the vacuum housing during operation of the reactive cleaning system, the backing pump (422) operable during operation of the reactive gas cleaning system.
- 25. The system for generating an ion beam according to any of the foregoing claims associated with an ion implantation apparatus, the apparatus constructed to transport ions

following the extraction electrode (405 or 220) to an implantation station (312, 314, FIG. 16) within a vacuum chamber (330).

- 26. The system for generating an ion beam of claim 25 including an isolation valve (425 or 200) for isolating the implantation station (312) from the ionization chamber (500) and the extraction electrode (405 or 220) during operation of the reactive gas cleaning system.
- 27. The system for generating an ion beam of any of the foregoing claims in which the ion source is constructed and adapted to generate dopant ions for semiconductor processing, and the reactive gas cleaning system (455, 430) is adapted to deliver fluorine, F, or chlorine, Cl, ions to the ionization chamber (500) or the extraction electrode (405 or 220) for cleaning deposits from a surface.
- 28. The system for generating an ion beam of any of the foregoing claims in which said ion source (400 or 10) is adapted to be temperature controlled to a given temperature.
- 29. The system for generating an ion beam of any of the foregoing claims in which said ion source (400 or 10) is adapted to generate a boron-containing ion beam.
- 30. The system for generating an ion beam of claim 29 adapted to generate said boron-containing ion beam by feeding vaporized borohydride material into said ion source (400 or 10).
- 31. The system for generating an ion beam of claim 30 in which said borohydride material is decaborane, $B_{10}H_{14}$.
- 32. The system for generating an ion beam of claim 30 in which said borohydride material is octadecaborane, $B_{12}H_{22}$.
- 33. The system for generating an ion beam of any of the claims 1-28 in which said ion source (400 or 10) is adapted to generate arsenic-containing ion beams.

34. The system for generating an ion beam of any of the claims 1-28 in which said ion source (400 or 10) is adapted to generate phosphorus-containing ion beams.

- 35. The system for generating an ion beam of any of the foregoing claims in which the ionization chamber (500) of said ion source comprises aluminum.
- 36. The system for generating an ion beam of any of the foregoing claims in which the ionization chamber (500) of said ion source or said extraction electrode comprises a material resistant to attack by halogen gases such as fluorine, F.
- 37. A method of in-situ cleaning with the system of any of the foregoing claims, or of an ion source associated with an ion implanter, in which reactive halogen gas is flowed into an ion source (400 or 10) while the ion source is de-energized and under vacuum.
- 38. The method of claim 37 in which the reactive halogen gas is fluorine, F.
- 39. The method of claim 37 in which the reactive halogen gas is chlorine, Cl.
- 40. The method of claim 37 in which the fluorine gas is introduced into the ion source from a remote plasma source.
- The method of claim 40 in which the fluorine gas is produced in the remote plasma source by an NF₃ plasma.
- 42. The method of claim 40 in which the fluorine gas is produced in the remote plasma source by a C_3F_3 or CF_4 plasma.
- 43. The method of claim 37 in which the reactive halogen gas is ClF₃.
- 44. The method of any of the claims 37-43 in which the cleaning procedure is conducted to remove deposits after the ion source has ionized decaborane, $\rm B_{10}H_{14}$.

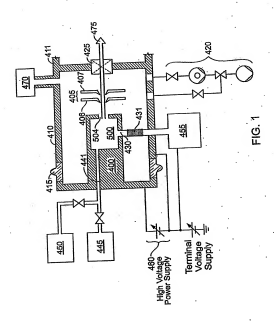
45. The method of any of the claims 37-43 in which the cleaning procedure is conducted to remove deposits after the ion source has ionized octadecaborane, B₁₈H₂₂.

- 46. The method of any of the claims 37-43 in which the cleaning procedure is conducted to remove deposits after the ion source has ionized arsenic-containing compounds, such as arsine, AsH₃, or elemental arsenic, As.
- 47. The method of any of the claims 37-43 in which the cleaning procedure is conducted to remove deposits after the ion source has ionized phosphorus-containing compounds, such as elemental phosphorus, P, or phosphine, PH₃.
- 48. The method of any of the claims 37-43 in which the cleaning procedure is conducted to remove deposits after the ion source has ionized antimony-containing compounds, such as trimethylantimony, Sb(CH₄)₃, or antimony pentaflouride, SbF₅.
- 49. The method of any of the claims 37-43 conducted for ion implanting in which the cleaning procedure is conducted for an ion source in situ in an ion implanter between changing ion source feed materials in order to implant a different ion species.
- 50. An ion implantation system having an ion source (400 or 10) and an extraction electrode (405 or 220) for extracting ions from said ion source, in which said electrode includes a heater constructed to maintain the electrode at an elevated temperature sufficient to substantially reduce condensation on said electrode of gases or vapors being ionized and products produced therefrom.
- The system of claim 50 in which said electrode comprises aluminum.
- 52. The system of claim 50 in which said electrode comprises of molybdenum.
- The system of claim 50, 51 or 52 in which said electrode is heated by radiative heaters (720, 730).

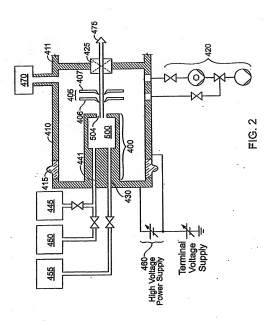
- 54. The system of claim 50, 51 or 52 in which said electrode is heated by resistive heaters (820).
- 55. The system of claim 53 or claim 54 in which the temperature of said electrode is controlled to a desired temperature.
- The system of claim 55 in which said temperature is between 150C and 250C.
- 57. The system of any of the claims 50-56 in which said electrode is periodically cleaned by exposure to reactive halogen-containing gas.
 - 58. A method of in-situ cleaning of an ion extraction electrode (405 or 220) of the system of any of the claims 1-36 or an ion extraction electrode (405 or 220) associated with an ion implanter, in which reactive halogen gas is flowed over the ion extraction electrode while said electrode is in situ and under vacuum.
 - 59. The method of claim 58 where the reactive halogen gas is fluorine, F.
 - 60. The method of claim 58 where the reactive halogen gas is chlorine, Cl.
 - 61. The method of claim 59 in which the fluorine gas is introduced from a remote plasma source (455) into a vacuum housing in which said extraction electrode resides.
 - 62. The method of claim 61 in which the fluorine gas is produced in the remote plasma source by a NF₃ plasma.
 - 63. The method of claim 61 in which the fluorine gas is produced in the remote plasma source by a C_3F_8 or CF_4 plasma.
 - 64. The method of claim 58 in which the reactive gas is CIP₃.

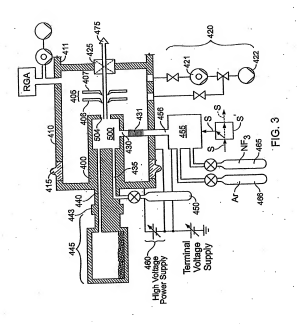
65. The method of any of the claims 58-64 in which the cleaning procedure is conducted to remove deposits after the ion source has ionized decaborane, $B_{10}H_{14}$.

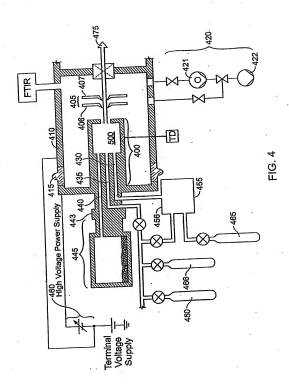
- 66. The method of any of the claims 58-64 in which the cleaning procedure is conducted to remove deposits after the ion source has ionized octadecaborane, B₁₈H₂₂.
- 67. The method of any of the claims 58-64 in which the cleaning procedure is conducted to remove deposits after the ion source has ionized arsenic-containing compounds, such as arsine, AsH₃, or elemental arsenic, As.
- 68. The method of any of the claims 58-64 in which the cleaning procedure is conducted to remove deposits after the ion source has ionized phosphorus-containing compounds, such as elemental phosphorus, P, or phosphine, PH₃.
- 69. The method of any of the claims 58-64 conducted in conjunction with ion implanting in which the cleaning procedure is conducted between changing ion source feed materials in order to implant a different ion species.

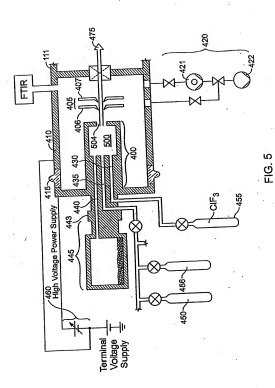


2/21-











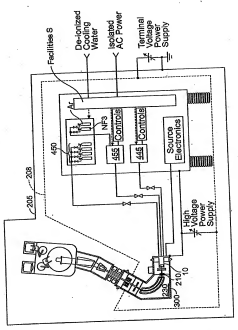


FIG. 6

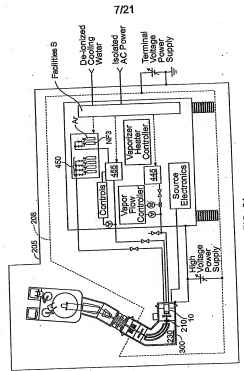
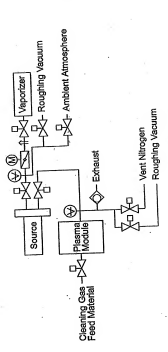


FIG. 6A



-1G. 6B

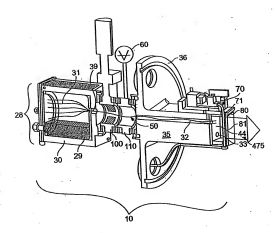


FIG. 7

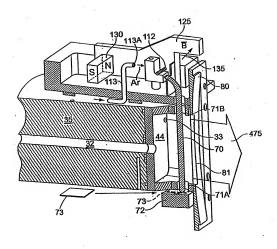
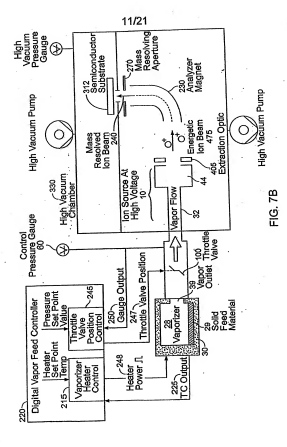


FIG. 7A



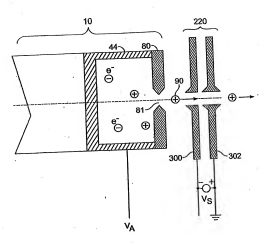
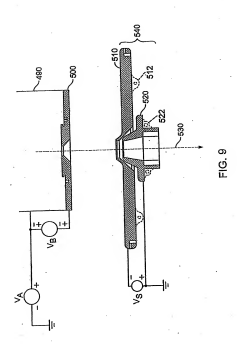
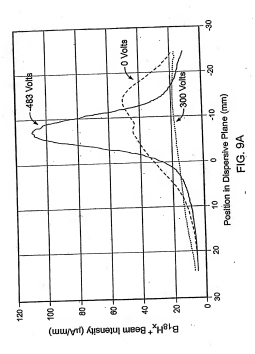
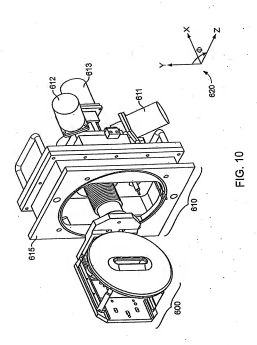


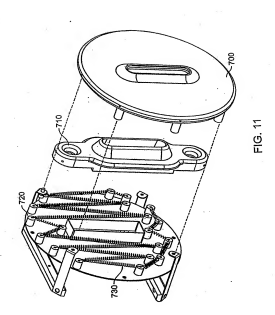
FIG. 8

13/21









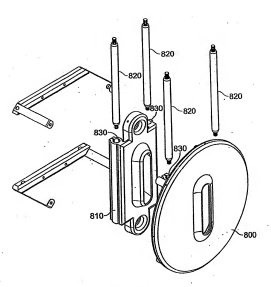
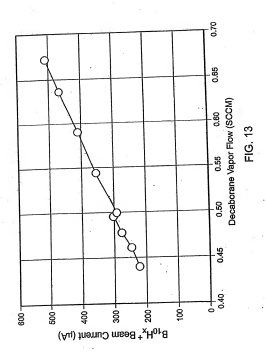


FIG. 12





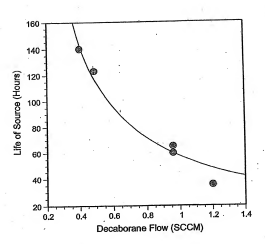
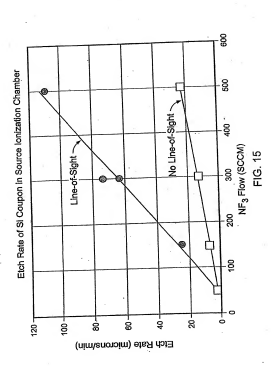


FIG. 14

20/21



21/21

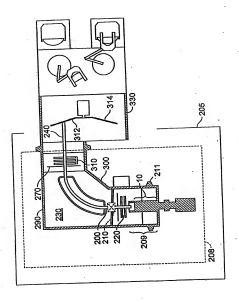


FIG. 16